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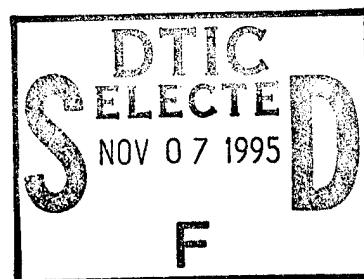


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PARALLEL SOLUTIONS OF TRIDIAGONAL AND PENTADIAGONAL SYSTEMS

by

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
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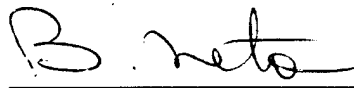
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
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

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PARALLEL SOLUTIONS OF TRIDIAGONAL AND PENTADIAGONAL SYSTEMS

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Abstract

We present an algorithm for the parallel solution of tridiagonal and pentadiagonal linear systems having nonzero elements at the top right and bottom left corners. Tridiagonal systems of this kind arise from the solution of two point boundary value problems with periodic boundary conditions. Pentadiagonal systems of this kind arise from e.g the approximation of the shallow water equations by the two-stage Galerkin method combined with a high accuracy compact approximation to the first derivative (Navon, 1983).

1. Introduction

In this paper, we develop an algorithm for the parallel solution of tridiagonal and pentadiagonal linear systems having nonzero elements at the top right and bottom left corners. This is a generalization of an algorithm due to Kowalik et al (1984) for tridiagonal systems. Such tridiagonal systems arise when approximating a class of two-point boundary value problems having periodic boundary conditions:

$$y''(t) = f(t, y(t)), \quad 0 \leq t \leq 1, \quad (1)$$

$$y(0) = y(1), \quad (2)$$

$$y'(0) = y'(1). \quad (3)$$

It was shown by Katti (1995) that this problem leads to the system

$$Ay = d \quad (4)$$

where the matrix A is of the form

$$\begin{pmatrix} a_1 & c_1 & 0 & 0 & \cdots & p_1 \\ b_2 & a_2 & c_2 & 0 & \cdots & 0 \\ 0 & b_3 & a_3 & c_3 & 0 & \cdots \\ \vdots & \vdots & \ddots & \ddots & \ddots & \\ 0 & \cdots & 0 & b_{N-1} & a_{N-1} & c_{N-1} \\ q_1 & \cdots & 0 & 0 & b_N & a_N \end{pmatrix} \quad (5)$$

and the right hand side d is

$$\begin{pmatrix} d_1 \\ d_2 \\ d_3 \\ \vdots \\ d_{N-1} \\ d_N \end{pmatrix}. \quad (6)$$

Pentadiagonal as well as tridiagonal systems appear when the two-stage Galerkin method combined with a high accuracy compact approximation to the first derivative is used for the approximation of the shallow water equations with

periodic boundary conditions (Navon, 1983). In this case, one has to solve a pentadiagonal system of the form

$$Bx = d \quad (7)$$

where the matrix B is given by

$$\begin{pmatrix} a_1 & c_1 & b_1 & 0 & 0 & 0 & \cdots & \cdots & p_1 & q_1 \\ r_2 & a_2 & c_2 & b_2 & 0 & 0 & \cdots & \cdots & 0 & q_2 \\ s_3 & r_3 & a_3 & c_3 & b_3 & 0 & \cdots & \cdots & 0 & 0 \\ 0 & s_4 & r_4 & a_4 & c_4 & b_4 & 0 & \cdots & 0 & 0 \\ \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots & \vdots \\ 0 & \cdots & 0 & s_{N-4} & r_{N-4} & a_{N-4} & c_{N-4} & b_{N-4} & 0 & 0 \\ 0 & 0 & \cdots & 0 & s_{N-3} & r_{N-3} & a_{N-3} & c_{N-3} & b_{N-3} & 0 \\ 0 & 0 & 0 & \cdots & 0 & s_{N-2} & r_{N-2} & a_{N-2} & c_{N-2} & b_{N-2} \\ u_1 & 0 & 0 & 0 & \cdots & 0 & s_{N-1} & r_{N-1} & a_{N-1} & c_{N-1} \\ v_1 & v_2 & 0 & 0 & \cdots & \cdots & 0 & s_N & r_N & a_N \end{pmatrix} \quad (8)$$

In the next section we describe the parallel algorithm for the direct solution of the tridiagonal system. The algorithm for the pentadiagonal system is described in section 3. Numerical experiments with both algorithms are reported in section 4. The two programs are attached as appendices.

2. Algorithm for Tridiagonal

In this section, we generalize the algorithm developed by Kowalik et al [1] for tridiagonal systems to the case where the matrix A is given by (5). We follow Kowalik in our description. Divide the N equations equally among the π processors (some may have 1 more equation than others). Let's assume for simplicity that each processor gets k equations. The first step is to eliminate b_j which are the elements below the diagonal. Each processor $1 \leq i \leq \pi$ eliminates b_j for $(i-1)k+2 \leq j \leq ik$.

For $j = (i-1)k+2, \dots, ik$

$f_{(i-1)k+1} \leftarrow b_{(i-1)k+1}$ only for $i \neq 1$

Compute the multiplier $m_j = \frac{b_j}{a_{j-1}}$ and update

$f_j \leftarrow f_j - m_j f_{j-1}$ only for $i \neq 1$

$a_j \leftarrow a_j - m_j c_{j-1}$

$p_j \leftarrow p_j - m_j p_{j-1}$ only for $i = 1$

$d_j \leftarrow d_j - m_j d_{j-1}$

This process creates the new elements f_j and p_j , as can be seen in (9) for the case $\pi = 3$.

$$\begin{pmatrix}
 a_1 & c_1 & \cdots & 0 & 0 & \cdots & \cdots & 0 & \cdots & \cdots & p_1 \\
 0 & a_2 & c_2 & & 0 & \cdots & \cdots & 0 & \cdots & \cdots & p_2 \\
 \vdots & & \ddots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 0 & 0 & \cdots & a_k & c_k & 0 & \cdots & 0 & \cdots & \cdots & p_k \\
 \hline
 0 & 0 & \cdots & f_{k+1} & a_{k+1} & c_{k+1} & \cdots & 0 & \cdots & \cdots & 0 \\
 \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & & \vdots & & & \vdots \\
 \vdots & \vdots & \vdots & \vdots & & \ddots & \ddots & \vdots & & & \vdots \\
 0 & 0 & \cdots & f_{2k} & 0 & \cdots & a_{2k} & c_{2k} & \cdots & \cdots & 0 \\
 \hline
 0 & 0 & \cdots & 0 & 0 & \cdots & f_{2k+1} & a_{2k+1} & c_{2k+1} & \cdots & 0 \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & & \vdots \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & & \ddots & \ddots & \vdots \\
 0 & 0 & \cdots & 0 & 0 & \cdots & f_{3k-1} & 0 & \cdots & a_{3k-1} & c_{3k-1} \\
 q_1 & 0 & \cdots & 0 & 0 & \cdots & f_{3k} & 0 & \cdots & \cdots & a_{3k}
 \end{pmatrix} \quad (9)$$

The second step is the elimination of c_j which are the elements above the diagonal. Each processor $1 \leq i \leq \pi$ eliminates c_j for $ik - 2 \leq j \leq (i - 1)k + 1$.

For $j = ik - 2, \dots, (i - 1)k + 1$

$g_{ik-1} \leftarrow c_{ik-1}$

Compute the multiplier $m_j = \frac{c_j}{a_{j+1}}$ and update

$f_j \leftarrow f_j - m_j f_{j+1}$ only for $i \neq 1$

$g_j \leftarrow g_j - m_j g_{j+1}$

$p_j \leftarrow p_j - m_j p_{j+1}$ only for $i = 1$

$d_j \leftarrow d_j - m_j d_{j+1}$

Note that for each processor i , the elements c_{ik} remain and can only be eliminated by passing the first row of processor $i + 1$. Upon completing these steps, the system becomes

$$\begin{pmatrix}
 a_1 & 0 & \cdots & g_1 & 0 & \cdots & \cdots & 0 & 0 & \cdots & \cdots & p_1 \\
 0 & a_2 & 0 & \vdots & 0 & \cdots & \cdots & 0 & 0 & \cdots & \cdots & p_2 \\
 \vdots & \vdots & \ddots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 \vdots & \vdots & & a_{k-1} & g_{k-1} & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 0 & 0 & \cdots & a_k & 0 & 0 & \cdots & g_k & 0 & \cdots & \cdots & p_k \\
 \hline
 0 & 0 & \cdots & f_{k+1} & a_{k+1} & 0 & \cdots & g_{k+1} & 0 & \cdots & \cdots & 0 \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 \vdots & \vdots & \vdots & \vdots & 0 & \cdots & a_{2k-1} & g_{2k-1} & \vdots & \vdots & \vdots & \vdots \\
 0 & 0 & \cdots & f_{2k} & 0 & \cdots & 0 & a_{2k} & 0 & \cdots & \cdots & g_{2k} \\
 \hline
 0 & 0 & \cdots & 0 & 0 & \cdots & f_{2k+1} & a_{2k+1} & 0 & \cdots & \cdots & g_{2k+1} \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\
 0 & 0 & \cdots & 0 & 0 & \cdots & f_{3k-1} & 0 & \cdots & a_{3k-1} & g_{3k-1} & \\
 q_1 & 0 & \cdots & 0 & 0 & \cdots & f_{3k} & 0 & \cdots & \cdots & a_{3k} &
 \end{pmatrix} \quad (10)$$

If we now take the first equation from processor 1 and the last equation from each processor, we end up with a similar tridiagonal system with only $\pi + 1$ equations:

$$\begin{pmatrix}
 a_1 & g_1 & 0 & p_1 \\
 f_k & a_k & g_k & 0 \\
 0 & f_{2k} & a_{2k} & g_{2k} \\
 q_1 & 0 & f_{3k} & a_{3k}
 \end{pmatrix} \quad (11)$$

where f_k is created upon elimination of p_k using the first row. In general, the matrix is

$$\begin{pmatrix}
 a_1 & g_1 & 0 & \cdots & \cdots & p_1 \\
 f_k & a_k & g_k & 0 & \cdots & 0 \\
 0 & f_{2k} & a_{2k} & g_{2k} & & \vdots \\
 \vdots & & \ddots & \ddots & \ddots & \\
 0 & \cdots & \cdots & f_{(\pi-1)k} & a_{(\pi-1)k} & g_{(\pi-1)k} \\
 q_1 & 0 & \cdots & 0 & f_{\pi k} & a_{\pi k}
 \end{pmatrix} \quad (12)$$

This system can be solved either serially or in parallel and is referred henceforth as the reduced system. The solution of this system is then broadcasted to all the processors. Upon receiving the solution of the reduced system, each processor i solves for the unknowns $x_{(i-1)k+1}, \dots, x_{ik-1}$ in parallel by a straightforward backward substitution. Note that for processor 1, the term x_1 need not be solved for since it is already known by virtue of the solution of the reduced system.

3. Algorithm for Pentadiagonal

In this section, we develop an algorithm for the pentadiagonal system (7). There are a few differences between this algorithm and the previous one. First, the reduced system contains the first two equations from the first processor and the last two equations from each processor. Thus the reduced system is of order $2(\pi + 1)$. Second, each processor i requires two equations from processor $i + 1$ to eliminate certain entries above the diagonal (c_j and b_j).

We now give the details of the algorithm, assuming again $\pi = 3$ processors each containing k equations. The first step is to eliminate the entries below the diagonal, namely s_j and r_j . Each processor $1 \leq i \leq \pi$ performs this task on the equations $(i - 1)k + 2 \leq j \leq ik - 1$.

For $j = (i - 1)k + 2, \dots, ik - 1$

$$f1_{(i-1)k+1} \leftarrow s_{(i-1)k+1} \quad \text{only for } i \neq 1$$

$$f2_{(i-1)k+1} \leftarrow r_{(i-1)k+1} \quad \text{only for } i \neq 1$$

$$f2_{(i-1)k+2} \leftarrow s_{(i-1)k+2} \quad \text{only for } i \neq 1$$

Compute the multiplier $m_j = \frac{r_j}{a_{j-1}}$ and update

$$f1_j \leftarrow f1_j - m_j f1_{j-1} \quad \text{only for } i \neq 1$$

$$f2_j \leftarrow f2_j - m_j f2_{j-1} \quad \text{only for } i \neq 1$$

$$a_j \leftarrow a_j - m_j c_{j-1}$$

$$c_j \leftarrow c_j - m_j b_{j-1}$$

$$p_j \leftarrow p_j - m_j p_{j-1} \quad \text{only for } i = 1$$

$$q_j \leftarrow q_j - m_j q_{j-1} \quad \text{only for } i = 1$$

$$d_j \leftarrow d_j - m_j d_{j-1}$$

Now compute the other multiplier $n_j = \frac{s_{j+1}}{a_{j-1}}$ and update

$$f1_{j+1} \leftarrow f1_{j+1} - n_j f1_{j-1} \quad \text{only for } i \neq 1$$

$$f2_{j+1} \leftarrow f2_{j+1} - n_j f2_{j-1} \quad \text{only for } i \neq 1$$

$$r_{j+1} \leftarrow r_{j+1} - n_j c_{j-1}$$

$$a_{j+1} \leftarrow a_{j+1} - n_j b_{j-1}$$

$$p_{j+1} \leftarrow p_{j+1} - n_j p_{j-1} \quad \text{only for } i = 1$$

$$q_{j+1} \leftarrow q_{j+1} - n_j q_{j-1} \quad \text{only for } i = 1$$

$$d_{j+1} \leftarrow d_{j+1} - n_j d_{j-1}$$

Note that for each processor i , the elements r_{ik} remain and are never eliminated. Because the semi-bandwidth of the matrix has been increased by one, then so does the number of equations that need to be passed between processors. The matrix B becomes

$$\begin{pmatrix}
 a_1 & c_1 & b_1 & \dots & 0 & \dots & \dots & \dots & 0 & \dots & p_1 & q_1 \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 0 & \dots & a_{k-1} & c_{k-1} & b_{k-1} & 0 & \dots & \dots & 0 & \dots & p_{k-1} & q_{k-1} \\
 0 & \dots & r_k & a_k & c_k & b_k & 0 & \dots & 0 & \dots & p_k & q_k \\
 \hline
 0 & \dots & f1_{k+1} & f2_{k+1} & a_{k+1} & c_{k+1} & b_{k+1} & \dots & 0 & \dots & \dots & 0 \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 0 & \dots & f1_{2k-1} & f2_{2k-1} & 0 & \dots & a_{2k-1} & c_{2k-1} & b_{2k-1} & 0 & \dots & 0 \\
 0 & \dots & f1_{2k} & f2_{2k} & 0 & \dots & r_{2k} & a_{2k} & c_{2k} & b_{2k} & 0 & \dots \\
 \hline
 0 & 0 & \dots & 0 & 0 & \dots & f1_{2k+1} & f2_{2k+1} & a_{2k+1} & c_{2k+1} & b_{2k+1} & \dots \\
 \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
 u_1 & 0 & \dots & 0 & 0 & \dots & f1_{3k-1} & f2_{3k-1} & 0 & \dots & a_{3k-1} & c_{3k-1} \\
 v_1 & v_2 & 0 & \dots & 0 & \dots & f1_{3k} & f2_{3k} & 0 & \dots & \dots & a_{3k}
 \end{pmatrix} \quad (13)$$

The second step is to eliminate the entries above the diagonal, namely c_j and b_j . Each processor $1 \leq i \leq \pi$ performs this task on the equations $ik - 3 \leq j \leq (i-1)k + 1$.

For $j = ik - 3, \dots, (i-1)k + 1$

$$g1_{ik-3} \leftarrow b_{ik-3}$$

$$g1_{ik-2} \leftarrow c_{ik-2}$$

$$g2_{ik-2} \leftarrow b_{ik-2}$$

Compute the multiplier $m_j = \frac{c_j}{a_{j+1}}$ and update

$$f1_j \leftarrow f1_j - m_j f1_{j+1} \quad \text{only for } i \neq 1$$

$$f2_j \leftarrow f2_j - m_j f2_{j+1} \quad \text{only for } i \neq 1$$

$$g1_j \leftarrow g1_j - m_j g1_{j+1}$$

$$g2_j \leftarrow g2_j - m_j g2_{j+1}$$

$$p_j \leftarrow p_j - m_j p_{j+1} \quad \text{only for } i = 1$$

$$q_j \leftarrow q_j - m_j q_{j+1} \quad \text{only for } i = 1$$

$$d_j \leftarrow d_j - m_j d_{j+1}$$

Now compute the other multiplier $n_j = \frac{b_{j-1}}{a_{j+1}}$ and update iff $j > (i-1)k + 1$

$$f1_{j-1} \leftarrow f1_{j-1} - n_j f1_{j+1} \quad \text{only for } i \neq 1$$

$$f2_{j-1} \leftarrow f2_{j-1} - n_j f2_{j+1} \quad \text{only for } i \neq 1$$

$$g1_{j-1} \leftarrow g1_{j-1} - n_j g1_{j+1}$$

$$g2_{j-1} \leftarrow g2_{j-1} - n_j g2_{j+1}$$

$$p_{j-1} \leftarrow p_{j-1} - n_j p_{j+1} \quad \text{only for } i = 1$$

$$q_{j-1} \leftarrow q_{j-1} - n_j q_{j+1} \quad \text{only for } i = 1$$

$$d_{j-1} \leftarrow d_{j-1} - n_j d_{j+1}$$

Note that this process does not eliminate the elements b_{ik-1} , b_{ik} , and c_{ik} for each processor i . Thus we have the following system

$$\begin{pmatrix}
a_1 & 0 & \cdots & g1_1 & g2_1 & 0 & \cdots & \cdots & \cdots & 0 & 0 & \cdots & \cdots & p_1 & q_1 \\
0 & a_2 & & g1_2 & g2_2 & 0 & \cdots & \cdots & \cdots & 0 & 0 & \cdots & \cdots & p_2 & q_2 \\
\vdots & & \ddots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
0 & 0 & \cdots & a_{k-1} & c_{k-1} & b_{k-1} & 0 & \cdots & \cdots & 0 & 0 & \cdots & \cdots & p_{k-1} & q_{k-1} \\
0 & 0 & \cdots & r_k & a_k & c_k & b_k & 0 & \cdots & 0 & 0 & \cdots & \cdots & p_k & q_k \\
\hline
0 & 0 & \cdots & f1_{k+1} & f2_{k+1} & a_{k+1} & 0 & \cdots & g1_{k+1} & g2_{k+1} & 0 & \cdots & \cdots & \cdots & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
\vdots & \vdots & \cdots & \vdots & \vdots & 0 & \cdots & \cdots & a_{2k-1} & c_{2k-1} & b_{2k-1} & 0 & \cdots & \cdots & 0 \\
0 & 0 & \cdots & f1_{2k} & f2_{2k} & 0 & \cdots & \cdots & r_{2k} & a_{2k} & c_{2k} & b_{2k} & 0 & \cdots & 0 \\
\hline
0 & 0 & \cdots & \cdots & 0 & 0 & \cdots & \cdots & f1_{2k+1} & f2_{2k+1} & a_{2k+1} & 0 & \cdots & g1_{2k+1} & g2_{2k+1} \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
b_{3k-1} & 0 & \cdots & \cdots & 0 & 0 & \cdots & \cdots & f1_{3k-1} & f2_{3k-1} & 0 & \cdots & \cdots & a_{3k-1} & c_{3k-1} \\
c_{3k} & b_{3k} & 0 & \cdots & 0 & 0 & \cdots & \cdots & f1_{3k} & f2_{3k} & 0 & \cdots & \cdots & r_{3k} & a_{3k}
\end{pmatrix} \quad (14)$$

To eliminate b_{ik-1} , b_{ik} and c_{ik} , the i^{th} processor requires the first two rows of processor $i + 1$. The elimination creates four new elements $g1_{ik-1}$, $g2_{ik-1}$, $g1_{ik}$, and $g2_{ik}$. By eliminating p_2 using row 1 and p_{k-1} , q_{k-1} , p_k , and q_k using rows 1 and 2 we then obtain the following reduced system

$$\begin{pmatrix}
a_1 & c_1 & g1_1 & g2_1 & 0 & 0 & p_1 & q_1 \\
r_2 & a_2 & g1_2 & g2_2 & 0 & 0 & 0 & q_2 \\
f1_{k-1} & f2_{k-1} & a_{k-1} & c_{k-1} & g1_{k-1} & g2_{k-1} & 0 & 0 \\
f1_k & f2_k & r_k & a_k & g1_k & g2_k & 0 & 0 \\
0 & 0 & f1_{2k-1} & f2_{2k-1} & a_{2k-1} & c_{2k-1} & g1_{2k-1} & g2_{2k-1} \\
0 & 0 & f1_{2k} & f2_{2k} & r_{2k} & a_{2k} & g1_{2k} & g2_{2k} \\
b_{3k-1} & 0 & 0 & 0 & f1_{3k-1} & f2_{3k-1} & a_{3k-1} & c_{3k-1} \\
c_{3k} & b_{3k} & 0 & 0 & f1_{3k} & f2_{3k} & r_{3k} & a_{3k}
\end{pmatrix} \quad (15)$$

Note that the elimination of p_2 only generates one new element r_2 , while the elimination of the other p 's and q 's creates the four new entries $f1_{k-1}$, $f2_{k-1}$, $f1_k$, and $f2_k$. Note that an additional term c_1 has appeared but this term is zero and is only included for illustrating the general structure of the reduced system. This matrix is a block tridiagonal system with nonzero blocks at the top right and bottom left corners, i.e. a block form of (11). This system can be solved serially by the block version of the solver used for (11). The solution is then broadcasted to all the processors. Upon receiving this solution, each processor i solves for the unknowns $x_{(i-1)k+2}, \dots, x_{ik-2}$ in parallel by a straightforward backward substitution. Note that for processor 1, the terms x_1 and x_2 need not be solved for since they are already known by virtue of the solution of the reduced system.

4. Numerical Experiments

In this section, we present the numerical experiments used to test the efficiency of the tridiagonal and pentadiagonal algorithms. For both algorithms, the number of processors π and equations per processor k are varied but the total number of equations remains fixed at $N = 10,000$. All runs are performed on a cluster of Sun4 workstations running PVM.

For the tridiagonal algorithm, the matrix shown in equation (5) is defined as

$$b_i = -1, \quad a_i = 2, \quad c_i = -1$$

and $d_i = 0$ for $i = 1, \dots, N$ with the additional conditions that $d_1 = -N$ and $d_N = N$. The corner terms in equation (5) are defined as $p_1 = b_1$ and $q_1 = c_N$. This system then has the exact solution $x_i = i$ for $i = 1, \dots, N$. Such a matrix system arises from a finite difference or finite element centered discretization of the one-dimensional Laplacian operator with periodic boundary conditions.

Table 1 shows the timing results obtained for the tridiagonal algorithm using averages based on five consecutive runs. Results are tabulated for $\pi = 2, 4, 8$ and 10 processors.

Number of Processors	Number of Equations Per Processor	Parallel Time (seconds)	Serial Time (seconds)	Speedup	Efficiency (%)
2	5000	0.10	0.11	1.1	55.0
4	2500	0.22	0.11	0.5	12.5
8	1250	0.22	0.11	0.5	6.3
10	1000	0.12	0.11	0.9	9.0

Table 1: Timings for the parallel and serial versions of the **tridiagonal** algorithm. Speedup is defined as the time ratio between the serial and parallel algorithm. Efficiency is the ratio of speedup to the number of processors.

The serial tridiagonal algorithm used for the purpose of computing speedup rates is a partitioning algorithm. The linear system (3) to be solved can be written as

$$Ax = b$$

which can then be partitioned as such

$$A(x^1 + x_n x^2) = b - A_{i,n} x_n$$

and this system can be decomposed into the two corresponding systems

$$\begin{aligned} Ax^1 &= b \\ Ax^2 &= -A_{i,n}. \end{aligned}$$

These two systems can now be solved via an LU decomposition. This algorithm is the periodic variant of the Thomas algorithm which is a very fast tridiagonal solver. Table 1 shows that the parallel version is competitive with the serial algorithm especially for $\pi = 2$ and $\pi = 10$ processors. For $\pi = 4$ and $\pi = 8$ processors the parallel algorithm is deficient relative to its serial counterpart. Nonetheless, these results demonstrate that no significant gains are made by parallelizing the tridiagonal solver.

For the pentadiagonal algorithm, the matrix shown in equation (8) is defined as

$$s_i = -1, \quad r_i = -1, \quad a_i = 4, \quad c_i = -1, \quad b_i = -1$$

and $d_i = 0$ for $i = 1, \dots, N$ with the additional conditions that $d_1 = -2N$, $d_2 = -N$, $d_{N-1} = N$ and $d_N = 2N$. The corner terms in equation (5) are defined as $p_1 = s_1$, $q_1 = r_1$, $q_2 = s_2$, $u_1 = b_{N-1}$, $v_1 = c_N$ and $v_2 = b_N$. This system then has the exact solution $x_i = i$ for $i = 1, \dots, N$. Such a matrix system arises from the finite difference centered discretization of the two-dimensional Laplacian operator with periodic boundary conditions.

Table 2 shows the timing results obtained for the pentadiagonal algorithm using averages based on five consecutive runs.

Number of Processors	Number of Equations Per Processor	Parallel Time (seconds)	Serial Time (seconds)	Speedup	Efficiency (%)
2	5000	0.11	0.22	2.0	100.0
4	2500	0.22	0.22	1.0	25.0
8	1250	0.22	0.22	1.0	12.5
10	1000	0.13	0.22	1.7	17.0

Table 2: Timings for the parallel and serial versions of the **pentadiagonal** algorithm. Speedup is defined as the time ratio between the serial and parallel algorithm. Efficiency is the ratio of speedup to the number of processors.

The serial pentadiagonal algorithm used for the purpose of computing speedup rates is the pentadiagonal version of the tridiagonal partitioning algorithm presented above. In this version the linear matrix system (8) can be written as

$$Bx = b$$

and partitioned as

$$B(x^1 + x_{n-1}x^2 + x_nx^3) = b - B_{i,n-1}x_{n-1} - B_{i,n}x_n$$

and this system can be decomposed into the three corresponding systems

$$Bx^1 = b$$

$$\begin{aligned} Bx^2 &= -B_{i,n-1} \\ Bx^3 &= -B_{i,n}. \end{aligned}$$

These three systems can now be solved via an LU decomposition. Table 2 shows that the parallel version does provide a significant gain over the serial algorithm for all of the number of processors studied. Once again, the major gains are achieved for $\pi = 2$ and $\pi = 10$ processors. In addition it is worth considering that the times for the parallel tridiagonal and pentadiagonal algorithms are almost identical. The communication is extremely detrimental to the overall performance of the tridiagonal algorithm, while for the pentadiagonal it begins to pay dividends. This study hints at the fact that as the bandwidth of the matrix is increased, the better the possible performance of this parallel algorithm versus its serial counterpart. The algorithm described is generalizable to any banded system. For a system with semi-bandwidth β , then β equations need to be passed between processors and the reduced system is of order $\beta(\pi + 1)$. Thus as the semi-bandwidth becomes sufficiently large, the majority of the communication time is spent on the actual transmission of information as opposed to the overhead incurred in communication calls.

5. Conclusions

An algorithm for tridiagonal and pentadiagonal matrices with nonzero elements at the top right and bottom left corners is presented. The algorithm is generalizable to higher banded systems but numerical studies are only performed for the tridiagonal and pentadiagonal cases. The numerical studies show that the communication overhead severely hurts the performance of the tridiagonal case. However, the results also show that gains are made in the pentadiagonal case and this trend points at the possibility of further gains when the bandwidth of the matrix is increased beyond the pentadiagonal case. The algorithm becomes a bit more cumbersome to implement but it does extend rather well to higher bandwidths. The number of equations that are required to be passed and the order of the reduced system increase at the rates β and $\beta(\pi + 1)$, respectively, where β is the semi-bandwidth of the matrix.

Acknowledgements

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Appendix A

In this appendix we give the tridiagonal solver. The first code is the parent (master) followed by the child (slave). The tridiagonal solver for the reduced system is also given.

```
*-----*
*This is the PARENT of a program which solves the
*tridiagonal system  $Ax = b$  with periodic boundary conditions using a
*Tridiagonal Decomposition Method
*as by B. Neta, C.P. Katti and F.X. Giraldo.
*Programmed by F.X. Giraldo on 7/95
*-----*

program parent3
include 'fpvm3.h'
include 'param.h'

                                !Global Arrays
dimension a(imax), b(imax), c(imax), d(imax), x(imax)
dimension a_p(nk), b_p(nk), c_p(nk), d_p(nk), xx(nk)
dimension fhat(nprocs+1), ahat(nprocs+1)
dimension ghat(nprocs+1), dhat(nprocs+1)
integer tids(16)
real taray(2)
character arch*8, nodename*10

npoin=nprocs*nk
do i=1,npoin                                !Construct Linear System
    b(i)=-1
    a(i)=4
    c(i)=-1
    d(i)=4 + 2*(i-2)
end do
d(1)=-npoin + 2
d(npoin)=3*npoin

                                !Start PVM
call pvmfmytid( mytid )
call pvmfparent( iptid )
nodename='child3.exe'
arch='*'

                                !Spawn Processors and distribute
                                !the Linear System among processors
do i=1,nprocs
    call pvmfspawn(nodename,pvmdefault,arch,1,tids(i),ierr)
```

```

write(*,'(" Spawning Processor tids ",i2,1x,i10)')i,tids(i)
if (ierr.ne.1) then
    write(*,'("ierr = ",i3)')ierr
    write(*,'("Error! Could not spawn process # ",i3)')i
    call pvmfexit(ierr)
    stop
endif
do j=1,nk
    b_p(j)=b( (i-1)*nk + j )
    a_p(j)=a( (i-1)*nk + j )
    c_p(j)=c( (i-1)*nk + j )
    d_p(j)=d( (i-1)*nk + j )
end do

msgtype=1
call pvmfinit send(pvmdefault,info)
call pvmfpack(integer4,i,1,1,info)
call pvmfpack(real4,b_p,nk,1,info)
call pvmfpack(real4,a_p,nk,1,info)
call pvmfpack(real4,c_p,nk,1,info)
call pvmfpack(real4,d_p,nk,1,info)
call pvmfpack(real4,tids,i,1,info)
call pvmf send(tids(i),msgtype,info)
end do

!Broadcast TIDS to all Processors
msgtype=2
call pvmfinit send(pvmdefault,info)
call pvmfpack(integer4,tids,16,1,info)
call pvmf mcast(nprocs,tids,msgtype,info)
!Construct Arrow System

time1=dtim e(taray)
msgtype=4
do i=1,nprocs
    call pvmfrecv(-1,msgtype,info)
    call pvmfunpack(integer4,iprocs,1,1,info)
    if (iprocs.eq.1) then
        !For i=1,2
        call pvmfunpack(real4,p1,1,1,info)
        call pvmfunpack(real4,a1,1,1,info)
        call pvmfunpack(real4,g1,1,1,info)
        call pvmfunpack(real4,d1,1,1,info)
        call pvmfunpack(real4,f2,1,1,info)
        call pvmfunpack(real4,a2,1,1,info)
        call pvmfunpack(real4,g2,1,1,info)
        call pvmfunpack(real4,d2,1,1,info)
        fhat(1)=p1
    end if
end do

```

```

    ahat(1)=a1
    ghat(1)=g1
    dhat(1)=d1
    fhat(2)=f2
    ahat(2)=a2
    ghat(2)=g2
    dhat(2)=d2
else if (iprocs.gt.1.and.iprocs.lt.nprocs) then !For i=3,NPROCS
    call pvmlunpack(real4,ff,1,1,info)
    call pvmlunpack(real4,aa,1,1,info)
    call pvmlunpack(real4,gg,1,1,info)
    call pvmlunpack(real4,dd,1,1,info)
    fhat(iprocs+1)=ff
    ahat(iprocs+1)=aa
    ghat(iprocs+1)=gg
    dhat(iprocs+1)=dd
else if (iprocs.eq.nprocs) then !For i=NPROCS + 1
    call pvmlunpack(real4,ff,1,1,info)
    call pvmlunpack(real4,aa,1,1,info)
    call pvmlunpack(real4,cc,1,1,info)
    call pvmlunpack(real4,dd,1,1,info)
    fhat(nprocs+1)=ff
    ahat(nprocs+1)=aa
    ghat(nprocs+1)=cc
    dhat(nprocs+1)=dd
endif
end do

!Solve Tridiagonal Arrow System
call tridiag_periodic(dhat,fhat,ahat,ghat,nprocs+1)
!Store Output in corresponding vector

x(1)=dhat(1)
x(nk)=dhat(2)
do i=3,nprocs+1
    x((i-1)*nk)=dhat(i)
end do

!Back Substitute
!Broadcast TIDS to all Processors

msgtype=5
call pvmlinitlsend(pvmldefault,info)
call pvmlpack(real4,x,imax,1,info)
call pvmlmcast(nprocs,tids,msgtype,info)
!Receive Local Sol from Children

msgtype=6
do i=1,nprocs

```

```

call pvmfrecv(-1,msgtype,info)
call pvmfunpack(integer4,iprocs,1,1,info)
call pvmfunpack(real4,xx,nk,1,info)
print*, ' Receiving local solution from Processor= ',iprocs
if (iprocs.eq.1) then
  do j=2, nk - 1
    x(j)=xx(j)
  end do
else
  do j=1, nk - 1
    x((iprocs-1)*nk+j)=xx(j)
  end do
endif
end do

time2=dttime(taray)
write(*, ' (" Total time in seconds = ",e12.4)')taray(1)+taray(2)

write(*, ' (" Storing Values " )')
open(1,file='parent3.out')
do i=1,npoin
  write(1, '(i7,1x,e16.8)')i,x(i)
end do
close(1)

stop
end

```

```

*-----*
*This is the CHILD of a program which solves the
*tridiagonal system  $A x = b$  with periodic boundary conditions using a
*Tridiagonal Decomposition Method
*as by B. Neta, C.P. Katti and F.X. Giraldo.
*Programmed by F.X. Giraldo on 7/95
*-----*

```

```

program child3
include 'fpvm3.h'
include 'param.h'

!Global Arrays
dimension a(nk), b(nk), c(nk), d(nk), x(imax), xx(nk)
dimension f(nk), g(nk), p(nk)
integer tids(16)

!Start PVM

call pvmfmytid( mytid )
call pvmfparent( mtid )

```

```

!Receive Data from Parent
msgtype=1
call pvmfrecv(mtid,msgtype,info)
call pvmfunpack(integer4,iprocs,1,1,info)
call pvmfunpack(real4,b,nk,1,info)
call pvmfunpack(real4,a,nk,1,info)
call pvmfunpack(real4,c,nk,1,info)
call pvmfunpack(real4,d,nk,1,info)

!Receive Broadcast
msgtype=2
call pvmfrecv(mtid,msgtype,info)
call pvmfunpack(integer4,tids,16,1,info)

!Eliminate b's
if (iprocs.eq.1) then
  p(1)=b(1)
  do j=2, nk
    xl=b(j)/a(j-1)
    a(j)=a(j) - xl*c(j-1)
    p(j)=-xl*p(j-1)
    d(j)=d(j) - xl*d(j-1)
  end do
else
  f(1)=b(1)
  do j=2, nk
    xl=b(j)/a(j-1)
    f(j)=-xl*f(j-1)
    a(j)=a(j) - xl*c(j-1)
    d(j)=d(j) - xl*d(j-1)
  end do
endif

!Eliminate c's
if (iprocs.eq.1) then
  g(nk-1)=c(nk-1)
  do j=nk - 2, 1, -1
    xl=c(j)/a(j+1)
    g(j)=-xl*g(j+1)
    p(j)=p(j) - xl*p(j+1)
    d(j)=d(j) - xl*d(j+1)
  end do
else
  g(nk-1)=c(nk-1)
  do j=nk - 2, 1, -1
    xl=c(j)/a(j+1)
    f(j)=f(j) - xl*f(j+1)

```



```

    call pvmfpack(real4,d(1),1,1,info)
    call pvmfpack(real4,f(nk),1,1,info)
    call pvmfpack(real4,a(nk),1,1,info)
    call pvmfpack(real4,g(nk),1,1,info)
    call pvmfpack(real4,d(nk),1,1,info)
else if (iprocs.gt.1.and.iprocs.lt.nprocs) then  !For i=3,NPROCS
    call pvmfpack(real4,f(nk),1,1,info)
    call pvmfpack(real4,a(nk),1,1,info)
    call pvmfpack(real4,g(nk),1,1,info)
    call pvmfpack(real4,d(nk),1,1,info)
else if (iprocs.eq.nprocs) then  !For i=NPROCS + 1
    call pvmfpack(real4,f(nk),1,1,info)
    call pvmfpack(real4,a(nk),1,1,info)
    call pvmfpack(real4,c(nk),1,1,info)
    call pvmfpack(real4,d(nk),1,1,info)
endif
call pvmfsend(mtid,msgtype,info)

                                !Back Substitute
                                !Receive Arrow Solution

msgtype=5
call pvmfrecv(mtid,msgtype,info)
call pvmfunpack(real4,x,imax,1,info)
                                !Obtain Local Solution

if (iprocs.eq.1) then
    do j=2, nk - 1
        xx(j)=( d(j) - g(j)*x(nk) - p(j)*x(nk*nprocs) )/a(j)
    end do
else
    do j=1, nk - 1
        xx(j)=( d(j) - f(j)*x((iprocs-1)*nk) - g(j)*x(iprocs*nk) )
$      /a(j)
    end do
endif

                                !Send Local Sol to Parent

msgtype=6
call pvmfinit send(pvmdefault,info)
call pvmfpack(integer4,iprocs,1,1,info)
call pvmfpack(real4,xx,nk,1,info)
call pvmfsend(mtid,msgtype,info)

stop
end
end

```

```

*This is the program which solves the
*reduced tridiagonal system  $Ax = b$  with periodic boundary conditions
*Programmed by F.X. Giraldo on 7/95
*-----*

      subroutine tridiag_periodic(d,b,a,c,n)
      include 'param.h'
      dimension b(nprocs+1), a(nprocs+1), c(nprocs+1), d(nprocs+1)
      dimension gam2(nprocs+1)

      a(1)=1.0/a(1)
      gam2(1)=-b(1)*a(1)
      b(1)=d(1)*a(1)
      do i=2,n-1
         c(i-1)=c(i-1)*a(i-1)
         a(i)=1.0/( a(i) - b(i)*c(i-1) )
         gam2(i)=-b(i)*gam2(i-1)*a(i)
         b(i)=( d(i) - b(i)*b(i-1) )*a(i)
      end do
      gam2(n-1)=gam2(n-1) - c(n-1)*a(n-1)

      d(n-1)=b(n-1)
      a(n-1)=gam2(n-1)
      do i1=2,n-1
         i=n-i1
         i2=i+1
         d(i)=b(i) - c(i)*d(i2)
         a(i)=gam2(i) - c(i)*a(i2)
      end do

      i=n
      zaa=d(i) - c(i)*d(1) - b(i)*d(n-1)
      zaa=zaa/( a(i) + b(i)*a(n-1) + c(i)*a(1) )
      d(i)=zaa
      do i=1,n-1
         d(i)=d(i)+a(i)*zaa
      end do

      return
      end

```


Appendix B

In this appendix we give the pentadiagonal solver. The first code is the parent (master) followed by the child (slave). The pentadiagonal solver for the reduced system is also given.

```
*-----*
*This is the PARENT of a program which solves the
*pentadiagonal system  $Ax = b$  with periodic boundary conditions using a
*Pentadiagonal Decomposition Method
*as by B. Neta, C.P. Katti and F. X. Giraldo.
*Programmed by F.X. Giraldo on 7/95
*-----*

      program parent5
      include 'fpvm3.h'
      include 'param.h'
c      external timing_fgettod

                                     !Global Arrays
      dimension s(imax), r(imax), a(imax), b(imax), c(imax)
      dimension d(imax), x(imax)
      dimension s_p(nk), r_p(nk), a_p(nk), b_p(nk), c_p(nk)
      dimension d_p(nk), x_p(nk)
      dimension fihat(nhat), f2hat(nhat), rhat(nhat)
      dimension ahat(nhat), chat(nhat), gihat(nhat)
      dimension g2hat(nhat), dhat(nhat), xhat(nhat)
      integer tids(nprocs), itime1(2), itime2(2), itotal
      real taray(2)
      character arch*8, nodename*10

      ichild=0
      npoin=nprocs*nk
      do i=1,npoin                                     !Construct Linear System
          s(i)=1
          r(i)=1
          a(i)=4
          c(i)=1
          b(i)=1
          d(i)=8*i
      end do
      d(1)=2*npoin + 8
      d(2)=npoin + 16
      d(npoin-1)=7*npoin - 8
      d(npoin)=6*npoin
```

```

                                !Start PVM

call pvmfmytid( mytid )
call pvmfparent( iptid )
nodename='child5.exe'
arch='*'

                                !Spawn Processors and distribute
                                !the Linear System among processors

do i=1,nprocs
  call pvmfspawn(nodename,pvmdefault,arch,1,tids(i),ierr)
  write(*,(' Spawning Processor tids ",i2,1x,i10)')i,tids(i)
  if (ierr.ne.1) then
    write(*,('ierr = ",i3)')ierr
    write(*,('Error! Could not spawn process # ",i3)')i
    call pvmfexit(ierr)
    stop
  endif
  do j=1,nk
    s_p(j)=s( (i-1)*nk + j )
    r_p(j)=r( (i-1)*nk + j )
    a_p(j)=a( (i-1)*nk + j )
    c_p(j)=c( (i-1)*nk + j )
    b_p(j)=b( (i-1)*nk + j )
    d_p(j)=d( (i-1)*nk + j )
  end do

  msgtype=1
  call pvmfinit send(pvmdefault,info)
  call pvmfpack(integer4,i,1,1,info)
  call pvmfpack(real4,s_p,nk,1,info)
  call pvmfpack(real4,r_p,nk,1,info)
  call pvmfpack(real4,a_p,nk,1,info)
  call pvmfpack(real4,c_p,nk,1,info)
  call pvmfpack(real4,b_p,nk,1,info)
  call pvmfpack(real4,d_p,nk,1,info)
  call pvmf send(tids(i),msgtype,info)
end do

                                !Broadcast TIDS to all Processors
c   call timing_fgettod(%REF(itime1))
  msgtype=2
  call pvmfinit send(pvmdefault,info)
  call pvmfpack(integer4,tids,nprocs,1,info)
  call pvmfmcast(nprocs,tids,msgtype,info)
                                !Construct Arrow System
  time1=dttime(taray)

```

```

msgtype=4
do i=1,nprocs
  call pvmfrecv(-1,msgtype,info)
  call pvmfunpack(integer4,iprocs,1,1,info)
  if (iprocs.eq.1) then
    !For i=1,2
    call pvmfunpack(real4,p1_1,1,1,info)
    call pvmfunpack(real4,p2_1,1,1,info)
    call pvmfunpack(real4,a_1,1,1,info)
    call pvmfunpack(real4,c_1,1,1,info)
    call pvmfunpack(real4,g1_1,1,1,info)
    call pvmfunpack(real4,g2_1,1,1,info)
    call pvmfunpack(real4,d_1,1,1,info)
    call pvmfunpack(real4,p2_2,1,1,info)
    call pvmfunpack(real4,r_2,1,1,info)
    call pvmfunpack(real4,a_2,1,1,info)
    call pvmfunpack(real4,g1_2,1,1,info)
    call pvmfunpack(real4,g2_2,1,1,info)
    call pvmfunpack(real4,d_2,1,1,info)
    call pvmfunpack(real4,f1_3,1,1,info)
    call pvmfunpack(real4,f2_3,1,1,info)
    call pvmfunpack(real4,a_3,1,1,info)
    call pvmfunpack(real4,c_3,1,1,info)
    call pvmfunpack(real4,g1_3,1,1,info)
    call pvmfunpack(real4,g2_3,1,1,info)
    call pvmfunpack(real4,d_3,1,1,info)
    call pvmfunpack(real4,f1_4,1,1,info)
    call pvmfunpack(real4,f2_4,1,1,info)
    call pvmfunpack(real4,r_4,1,1,info)
    call pvmfunpack(real4,a_4,1,1,info)
    call pvmfunpack(real4,g1_4,1,1,info)
    call pvmfunpack(real4,g2_4,1,1,info)
    call pvmfunpack(real4,d_4,1,1,info)
    pihat=p1_1
    p2hat=p2_1
    ahat(1)=a_1
    chat(1)=c_1
    g1hat(1)=g1_1
    g2hat(1)=g2_1
    dhat(1)=d_1
    q2hat=p2_2
    rhat(2)=r_2
    ahat(2)=a_2
    g1hat(2)=g1_2
    g2hat(2)=g2_2
  end if
end do

```

```

        dhat(2)=d_2
        f1hat(2*iprocs+1)=f1_3
f2hat(2*iprocs+1)=f2_3
        ahat(2*iprocs+1)=a_3
        chat(2*iprocs+1)=c_3
        g1hat(2*iprocs+1)=g1_3
        g2hat(2*iprocs+1)=g2_3
        dhat(2*iprocs+1)=d_3
        f1hat(2*iprocs+2)=f1_4
        f2hat(2*iprocs+2)=f2_4
        rhat(2*iprocs+2)=r_4
        ahat(2*iprocs+2)=a_4
        g1hat(2*iprocs+2)=g1_4
        g2hat(2*iprocs+2)=g2_4
        dhat(2*iprocs+2)=d_4
    else if (iprocs.gt.1.and.iprocs.lt.nprocs) then !For i=3,NPROCS
        call pvmfunpack(real4,f1_1,1,1,info)
        call pvmfunpack(real4,f2_1,1,1,info)
        call pvmfunpack(real4,a_1,1,1,info)
        call pvmfunpack(real4,c_1,1,1,info)
        call pvmfunpack(real4,g1_1,1,1,info)
        call pvmfunpack(real4,g2_1,1,1,info)
        call pvmfunpack(real4,d_1,1,1,info)
        call pvmfunpack(real4,f1_2,1,1,info)
        call pvmfunpack(real4,f2_2,1,1,info)
        call pvmfunpack(real4,r_2,1,1,info)
        call pvmfunpack(real4,a_2,1,1,info)
        call pvmfunpack(real4,g1_2,1,1,info)
        call pvmfunpack(real4,g2_2,1,1,info)
        call pvmfunpack(real4,d_2,1,1,info)
        f1hat(2*iprocs+1)=f1_1
f2hat(2*iprocs+1)=f2_1
        ahat(2*iprocs+1)=a_1
        chat(2*iprocs+1)=c_1
        g1hat(2*iprocs+1)=g1_1
        g2hat(2*iprocs+1)=g2_1
        dhat(2*iprocs+1)=d_1
        f1hat(2*iprocs+2)=f1_2
        f2hat(2*iprocs+2)=f2_2
        rhat(2*iprocs+2)=r_2
        ahat(2*iprocs+2)=a_2
        g1hat(2*iprocs+2)=g1_2
        g2hat(2*iprocs+2)=g2_2
        dhat(2*iprocs+2)=d_2

```

```

else if (iprocs.eq.nprocs) then
    call pvmfunpack(real4,f1_1,1,1,info)
    call pvmfunpack(real4,f2_1,1,1,info)
    call pvmfunpack(real4,a_1,1,1,info)
    call pvmfunpack(real4,c_1,1,1,info)
    call pvmfunpack(real4,b_1,1,1,info)
    call pvmfunpack(real4,d_1,1,1,info)
    call pvmfunpack(real4,f1_2,1,1,info)
    call pvmfunpack(real4,f2_2,1,1,info)
    call pvmfunpack(real4,r_2,1,1,info)
    call pvmfunpack(real4,a_2,1,1,info)
    call pvmfunpack(real4,c_2,1,1,info)
    call pvmfunpack(real4,b_2,1,1,info)
    call pvmfunpack(real4,d_2,1,1,info)
    f1hat(2*iprocs+1)=f1_1
    f2hat(2*iprocs+1)=f2_1
    ahat(2*iprocs+1)=a_1
    chat(2*iprocs+1)=c_1
    v1hat=b_1
    dhat(2*iprocs+1)=d_1
    f1hat(2*iprocs+2)=f1_2
    f2hat(2*iprocs+2)=f2_2
    rhat(2*iprocs+2)=r_2
    ahat(2*iprocs+2)=a_2
    u1hat=c_2
    u2hat=b_2
    dhat(2*iprocs+2)=d_2
endif
end do

                                !Solve Tridiagonal Arrow System
call pentadiag_periodic(f1hat,f2hat,rhat,ahat,chat,g1hat,
$    g2hat,p1hat,p2hat,q2hat,v1hat,u1hat,u2hat,dhat,xhat)

                                !Store Output in corresponding vector

x(1)=xhat(1)
x(2)=xhat(2)
do i=1,nprocs
    x(i*nk-1)=xhat(2*i+1)
    x(i*nk)=xhat(2*i+2)
end do

                                !Back Substitute
                                !Broadcast TIDS to all Processors

msgtype=5
call pvmfinitsend(pvmdefault,info)

```

```

call pvmfpack(real4,x,imax,1,info)
call pvmfmcast(nprocs,tids,msgtype,info)
                                !Receive Local Solution from Children
msgtype=6
do i=1,nprocs
  call pvmfrecv(-1,msgtype,info)
  call pvmfunpack(integer4,iprocs,1,1,info)
  call pvmfunpack(real4,x_p,nk,1,info)
  call pvmfunpack(integer4,itime,1,1,info)
  ichild=ichild + itime
  print*, ' Receiving solution Processor = ',iprocs
  if (iprocs.eq.1) then
    do j=3, nk - 2
      x(j)=x_p(j)
    end do
  else
    do j=1, nk - 2
      x((iprocs-1)*nk+j)=x_p(j)
    end do
  endif
end do

c   call timing_fgettod(%REF(itime2))
c   iparent=(itime2(1)-itime1(1))*1000000 + itime2(2)-itime1(2)
c   print*, ' Children Time in useconds= ',ichild
c   print*, ' Parent Time in useconds  = ',iparent
c   print*, ' Total Time in useconds   = ',ichild+iparent

time2=dttime(taray)
write(*, '(' Total time in seconds = ",e12.4)')taray(1)+taray(2)

write(*, '(' Storing Values " ')')
open(1,file='parent5.out')
do i=1,npoin
  write(1, '(i7,1x,e12.6)')i,x(i)
end do
close(1)

stop
end

*-----*
*This is the CHILD of a program which solves the
*pentadiagonal system  $A x = b$  with periodic boundary conditions using a
*Pentadiagonal Decomposition Method

```

*as by B. Neta, C.P. Katti and F. X. Giraldo

*Programmed by F.X. Giraldo on 7/95

```
*-----*
      program child5
      include 'fpvm3.h'
      include 'param.h'
c      external timing_fgettod

                                     !Global Arrays
      dimension s(nk), r(nk), a(nk), c(nk), b(nk), d(nk)
      dimension xg(imax), x(nk)
      dimension f1(nk), f2(nk), g1(nk), g2(nk), p1(nk), p2(nk)
      integer tids(nprocs), itime1(2), itime2(2), itotal
                                     !Start PVM

      call pvmfmytid( mytid )
      call pvmfparent( mtid )

                                     !Receive Data from Parent

      msgtype=1
      call pvmfrecv(mtid,msgtype,info)
      call pvmfunpack(integer4,iprocs,1,1,info)
      call pvmfunpack(real4,s,nk,1,info)
      call pvmfunpack(real4,r,nk,1,info)
      call pvmfunpack(real4,a,nk,1,info)
      call pvmfunpack(real4,c,nk,1,info)
      call pvmfunpack(real4,b,nk,1,info)
      call pvmfunpack(real4,d,nk,1,info)

                                     !Receive Broadcast
c      call timing_fgettod(%REF(itime1))
      msgtype=2
      call pvmfrecv(mtid,msgtype,info)
      call pvmfunpack(integer4,tids,nprocs,1,info)
                                     !Eliminate R and S

      if (iprocs.eq.1) then
        p1(1)=s(1)
        p2(1)=r(1)
      p2(2)=s(2)
        do j=2,nk-1
          x1=r(j)/a(j-1)
          a(j)=a(j) - x1*c(j-1)
          c(j)=c(j) - x1*b(j-1)
          p1(j)=p1(j) - x1*p1(j-1)
          p2(j)=p2(j) - x1*p2(j-1)
          d(j)=d(j) - x1*d(j-1)
          x1=s(j+1)/a(j-1)
          r(j+1)=r(j+1) - x1*c(j-1)
        end do
      end if
    end program child5
```

```

        a(j+1)=a(j+1) - x1*b(j-1)
        p1(j+1)=p1(j+1) - x1*p1(j-1)
        p2(j+1)=p2(j+1) - x1*p2(j-1)
        d(j+1)=d(j+1) - x1*d(j-1)
    end do
    else
f1(1)=s(1)
f2(1)=r(1)
f2(2)=s(2)
        do j=2,nk-1
            x1=r(j)/a(j-1)
            f1(j)=f1(j) - x1*f1(j-1)
            f2(j)=f2(j) - x1*f2(j-1)
            a(j)=a(j) - x1*c(j-1)
            c(j)=c(j) - x1*b(j-1)
            d(j)=d(j) - x1*d(j-1)
            x1=s(j+1)/a(j-1)
            f1(j+1)=f1(j+1) - x1*f1(j-1)
            f2(j+1)=f2(j+1) - x1*f2(j-1)
            r(j+1)=r(j+1) - x1*c(j-1)
            a(j+1)=a(j+1) - x1*b(j-1)
            d(j+1)=d(j+1) - x1*d(j-1)
        end do
    endif
!Eliminate C and B
g1(nk-3)=b(nk-3)
g1(nk-2)=c(nk-2)
g2(nk-2)=b(nk-2)
if (iprocs.eq.1) then
    do j=nk - 3,1, -1
        x1=c(j)/a(j+1)
        g1(j)=g1(j) - x1*g1(j+1)
        g2(j)=g2(j) - x1*g2(j+1)
        p1(j)=p1(j) - x1*p1(j+1)
        p2(j)=p2(j) - x1*p2(j+1)
        d(j)=d(j) - x1*d(j+1)
    if (j.gt.1) then
        x1=b(j-1)/a(j+1)
        g1(j-1)=g1(j-1) - x1*g1(j+1)
        g2(j-1)=g2(j-1) - x1*g2(j+1)
        p1(j-1)=p1(j-1) - x1*p1(j+1)
        p2(j-1)=p2(j-1) - x1*p2(j+1)
        d(j-1)=d(j-1) - x1*d(j+1)
    endif
endif

```



```

        end do
    else
        do j=nk - 3,1, -1
            x1=c(j)/a(j+1)
            f1(j)=f1(j) - x1*f1(j+1)
            f2(j)=f2(j) - x1*f2(j+1)
            g1(j)=g1(j) - x1*g1(j+1)
            g2(j)=g2(j) - x1*g2(j+1)
            d(j)=d(j) - x1*d(j+1)
            if (j.gt.1) then
                x1=b(j-1)/a(j+1)
                f1(j-1)=f1(j-1) - x1*f1(j+1)
                f2(j-1)=f2(j-1) - x1*f2(j+1)
                g1(j-1)=g1(j-1) -x1*g1(j+1)
                g2(j-1)=g2(j-1) -x1*g2(j+1)
                d(j-1)=d(j-1) - x1*d(j+1)
            endif
        end do
    endif

    !Send variables from i to i-1
    if (iprocs.gt.1) then
        msgtype=3
        call pvmfinit send(pvmdefault,info)
        call pvmfpack(real4,f1(1),1,1,info)
        call pvmfpack(real4,f2(1),1,1,info)
        call pvmfpack(real4,a(1),1,1,info)
        call pvmfpack(real4,g1(1),1,1,info)
        call pvmfpack(real4,g2(1),1,1,info)
        call pvmfpack(real4,d(1),1,1,info)
        call pvmfpack(real4,f1(2),1,1,info)
        call pvmfpack(real4,f2(2),1,1,info)
        call pvmfpack(real4,a(2),1,1,info)
        call pvmfpack(real4,g1(2),1,1,info)
        call pvmfpack(real4,g2(2),1,1,info)
        call pvmfpack(real4,d(2),1,1,info)
        call pvmf send(tids(iprocs-1),msgtype,info)
    endif

    !Receive variables from i+1 to i
    if (iprocs.lt.nprocs) then
        msgtype=3
        call pvmfrecv(tids(iprocs+1),msgtype,info)
        call pvmfunpack(real4,f1_1,1,1,info)
        call pvmfunpack(real4,f2_1,1,1,info)
        call pvmfunpack(real4,a_1,1,1,info)
    endif

```

```

call pvmpunpack(real4,g1_1,1,1,info)
call pvmpunpack(real4,g2_1,1,1,info)
call pvmpunpack(real4,d_1,1,1,info)
call pvmpunpack(real4,f1_2,1,1,info)
call pvmpunpack(real4,f2_2,1,1,info)
call pvmpunpack(real4,a_2,1,1,info)
call pvmpunpack(real4,g1_2,1,1,info)
call pvmpunpack(real4,g2_2,1,1,info)
call pvmpunpack(real4,d_2,1,1,info)
!Eliminate c_(nk), b_(nk) and b_(nk-1)

xl=c(nk)/a_1
r(nk)=r(nk) - xl*f1_1
a(nk)=a(nk) - xl*f2_1
g1(nk)=-xl*g1_1
g2(nk)=-xl*g2_1
d(nk)=d(nk) - xl*d_1
xl=b(nk)/a_2
r(nk)=r(nk) - xl*f1_2
a(nk)=a(nk) - xl*f2_2
g1(nk)=g1(nk) - xl*g1_2
g2(nk)=g2(nk) - xl*g2_2
d(nk)=d(nk) - xl*d_2
xl=b(nk-1)/a_1
a(nk-1)=a(nk-1) - xl*f1_1
c(nk-1)=c(nk-1) - xl*f2_1
g1(nk-1)=-xl*g1_1
g2(nk-1)=-xl*g2_1
d(nk-1)=d(nk-1) - xl*d_1
endif

!For Processor 1 Only

if (iprocs.eq.1) then
c(1)=0.0

!Eliminate P1_2

xl=p1(2)/p1(1)
r(2)=-xl*a(1)
g1(2)=g1(2) - xl*g1(1)
g2(2)=g2(2) - xl*g2(1)
p2(2)=p2(2) - xl*p2(1)
d(2)=d(2) - xl*d(1)

!Eliminate P1_nk-1

xl=p1(nk-1)/p1(1)
f1(nk-1)=-xl*a(1)
a(nk-1)=a(nk-1) - xl*g1(1)
c(nk-1)=c(nk-1) - xl*g2(1)

```

```

p2(nk-1)=p2(nk-1) - x1*p2(1)
d(nk-1)=d(nk-1) - x1*d(1)
                                !Eliminate P1_nk

x1=p1(nk)/p1(1)
f1(nk)=-x1*a(1)
r(nk)=r(nk) - x1*g1(1)
a(nk)=a(nk) - x1*g2(1)
p2(nk)=p2(nk) - x1*p2(1)
d(nk)=d(nk) - x1*d(1)
                                !Eliminate P2_nk-1

x1=p2(nk-1)/p2(2)
f1(nk-1)=f1(nk-1) - x1*r(2)
f2(nk-1)=-x1*a(2)
a(nk-1)=a(nk-1) - x1*g1(2)
c(nk-1)=c(nk-1) - x1*g2(2)
d(nk-1)=d(nk-1) - x1*d(2)
                                !Eliminate P2_nk

x1=p2(nk)/p2(2)
f1(nk)=f1(nk) - x1*r(2)
f2(nk)=-x1*a(2)
r(nk)=r(nk) - x1*g1(2)
a(nk)=a(nk) - x1*g2(2)
d(nk)=d(nk) - x1*d(2)
endif

                                !Construct Arrow System
msgtype=4
call pvmfinit(send(pvmdefault,info)
call pvmfpack(integer4,iprocs,1,1,info)
if (iprocs.eq.1) then
                                !For i=1,2
    call pvmfpack(real4,p1(1),1,1,info)
    call pvmfpack(real4,p2(1),1,1,info)
    call pvmfpack(real4,a(1),1,1,info)
    call pvmfpack(real4,c(1),1,1,info)
    call pvmfpack(real4,g1(1),1,1,info)
    call pvmfpack(real4,g2(1),1,1,info)
    call pvmfpack(real4,d(1),1,1,info)
    call pvmfpack(real4,p2(2),1,1,info)
    call pvmfpack(real4,r(2),1,1,info)
    call pvmfpack(real4,a(2),1,1,info)
    call pvmfpack(real4,g1(2),1,1,info)
    call pvmfpack(real4,g2(2),1,1,info)
    call pvmfpack(real4,d(2),1,1,info)
    call pvmfpack(real4,f1(nk-1),1,1,info)
    call pvmfpack(real4,f2(nk-1),1,1,info)

```

```

call pvmfpack(real4,a(nk-1),1,1,info)
call pvmfpack(real4,c(nk-1),1,1,info)
call pvmfpack(real4,g1(nk-1),1,1,info)
call pvmfpack(real4,g2(nk-1),1,1,info)
call pvmfpack(real4,d(nk-1),1,1,info)
call pvmfpack(real4,f1(nk),1,1,info)
call pvmfpack(real4,f2(nk),1,1,info)
call pvmfpack(real4,r(nk),1,1,info)
call pvmfpack(real4,a(nk),1,1,info)
call pvmfpack(real4,g1(nk),1,1,info)
call pvmfpack(real4,g2(nk),1,1,info)
call pvmfpack(real4,d(nk),1,1,info)
else if (iprocs.gt.1.and.iprocs.lt.nprocs) then !For i=3,NPROCS
call pvmfpack(real4,f1(nk-1),1,1,info)
call pvmfpack(real4,f2(nk-1),1,1,info)
call pvmfpack(real4,a(nk-1),1,1,info)
call pvmfpack(real4,c(nk-1),1,1,info)
call pvmfpack(real4,g1(nk-1),1,1,info)
call pvmfpack(real4,g2(nk-1),1,1,info)
call pvmfpack(real4,d(nk-1),1,1,info)
call pvmfpack(real4,f1(nk),1,1,info)
call pvmfpack(real4,f2(nk),1,1,info)
call pvmfpack(real4,r(nk),1,1,info)
call pvmfpack(real4,a(nk),1,1,info)
call pvmfpack(real4,g1(nk),1,1,info)
call pvmfpack(real4,g2(nk),1,1,info)
call pvmfpack(real4,d(nk),1,1,info)
else if (iprocs.eq.nprocs) then !For i=NPROCS+1
call pvmfpack(real4,f1(nk-1),1,1,info)
call pvmfpack(real4,f2(nk-1),1,1,info)
call pvmfpack(real4,a(nk-1),1,1,info)
call pvmfpack(real4,c(nk-1),1,1,info)
call pvmfpack(real4,b(nk-1),1,1,info)
call pvmfpack(real4,d(nk-1),1,1,info)
call pvmfpack(real4,f1(nk),1,1,info)
call pvmfpack(real4,f2(nk),1,1,info)
call pvmfpack(real4,r(nk),1,1,info)
call pvmfpack(real4,a(nk),1,1,info)
call pvmfpack(real4,c(nk),1,1,info)
call pvmfpack(real4,b(nk),1,1,info)
call pvmfpack(real4,d(nk),1,1,info)
endif
call pvmfsend(mtld,msgtype,info)
!Back Substitute

```

```

                                !Receive Arrow Solution
msgtype=5
call pvmfrecv(mtid,msgtype,info)
call pvmfunpack(real4,xg,imax,1,info)
                                !Obtain Local Solution
if (iprocs.eq.1) then
  do j=3, nk - 2
    x(j)=( d(j) - g1(j)*xg(nk-1) - g2(j)*xg(nk)
$         - p1(j)*xg(nk*nprocs-1) - p2(j)*xg(nk*nprocs) )/a(j)
  end do
else
  do j=1, nk - 2
    x(j)=( d(j) - f1(j)*xg((iprocs-1)*nk-1)
$         - f2(j)*xg((iprocs-1)*nk) - g1(j)*xg(iprocs*nk-1)
$         - g2(j)*xg(iprocs*nk) )/a(j)
  end do
endif
c   call timing_fgettod(%REF(itime2))
c   itotal=(itime2(1)-itime1(1))*1000000 + itime2(2)-itime1(2)
                                !Send Local Solution to Parent
msgtype=6
call pvmfinitssend(pvmdefault,info)
call pvmfpack(integer4,iprocs,1,1,info)
call pvmfpack(real4,x,nk,1,info)
call pvmfpack(integer4,itotal,1,1,info)
call pvmfssend(mtid,msgtype,info)

stop
end

*-----*
*This program solves a
*smaller pentadiagonal system  $A x = b$  with periodic boundary conditions
*using a matrix partitioning approach.
*Written on 7/95, by F. X. Giraldo
*-----*

subroutine pentadiag_periodic(f1hat,f2hat,rhat,ahat,chat,g1hat,
$      g2hat,p1hat,p2hat,q2hat,v1hat,u1hat,u2hat,dhat,xhat)
include 'param.h'
dimension f1hat(nhat), f2hat(nhat), rhat(nhat), ahat(nhat)
dimension chat(nhat), g1hat(nhat), g2hat(nhat), dhat(nhat)
dimension xhat(nhat), d2hat(nhat), d3hat(nhat)
dimension x1hat(nhat), x2hat(nhat), x3hat(nhat)
                                !Forward Reduction
                                !Forward Reduction

```

```

d2hat(1)=-p1hat
d2hat(nhat-3)=-g1hat(nhat-3)
d2hat(nhat-2)=-g1hat(nhat-2)
d3hat(1)=-p2hat
d3hat(2)=-q2hat
d3hat(nhat-3)=-g2hat(nhat-3)
d3hat(nhat-2)=-g2hat(nhat-2)

do i=1,nhat-5,2

                                !Eliminate rhat_i+1
    xl=rhat(i+1)/ahat(i)
    ahat(i+1)=ahat(i+1) - xl*chat(i)
    g1hat(i+1)=g1hat(i+1) - xl*g1hat(i)
    g2hat(i+1)=g2hat(i+1) - xl*g2hat(i)
    dhat(i+1)=dhat(i+1) - xl*dhat(i)
    d2hat(i+1)=d2hat(i+1) - xl*d2hat(i)
    d3hat(i+1)=d3hat(i+1) - xl*d3hat(i)

                                !Eliminate f1hat_i+2
    xl=f1hat(i+2)/ahat(i)
    f2hat(i+2)=f2hat(i+2) - xl*chat(i)
    ahat(i+2)=ahat(i+2) - xl*g1hat(i)
    chat(i+2)=chat(i+2) - xl*g2hat(i)
    dhat(i+2)=dhat(i+2) - xl*dhat(i)
    d2hat(i+2)=d2hat(i+2) - xl*d2hat(i)
    d3hat(i+2)=d3hat(i+2) - xl*d3hat(i)

                                !Eliminate f1hat_i+3
    xl=f1hat(i+3)/ahat(i)
    f2hat(i+3)=f2hat(i+3) - xl*chat(i)
    rhat(i+3)=rhat(i+3) - xl*g1hat(i)
    ahat(i+3)=ahat(i+3) - xl*g2hat(i)
    dhat(i+3)=dhat(i+3) - xl*dhat(i)
    d2hat(i+3)=d2hat(i+3) - xl*d2hat(i)
    d3hat(i+3)=d3hat(i+3) - xl*d3hat(i)

                                !Eliminate f2hat_i+2
    xl=f2hat(i+2)/ahat(i+1)
    ahat(i+2)=ahat(i+2) - xl*g1hat(i+1)
    chat(i+2)=chat(i+2) - xl*g2hat(i+1)
    dhat(i+2)=dhat(i+2) - xl*dhat(i+1)
    d2hat(i+2)=d2hat(i+2) - xl*d2hat(i+1)
    d3hat(i+2)=d3hat(i+2) - xl*d3hat(i+1)

                                !Eliminate f2hat_i+3
    xl=f2hat(i+3)/ahat(i+1)
    rhat(i+3)=rhat(i+3) - xl*g1hat(i+1)
    ahat(i+3)=ahat(i+3) - xl*g2hat(i+1)

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dhat(i+3)=dhat(i+3) - x1*dhat(i+1)
d2hat(i+3)=d2hat(i+3) - x1*d2hat(i+1)
d3hat(i+3)=d3hat(i+3) - x1*d3hat(i+1)
end do

!Eliminate rhat_nhat-3

i=nhat - 3
x1=rhat(i+1)/ahat(i)
ahat(i+1)=ahat(i+1) - x1*chat(i)
g1hat(i+1)=g1hat(i+1) - x1*g1hat(i)
g2hat(i+1)=g2hat(i+1) - x1*g2hat(i)
dhat(i+1)=dhat(i+1) - x1*dhat(i)
d2hat(i+1)=d2hat(i+1) - x1*d2hat(i)
d3hat(i+1)=d3hat(i+1) - x1*d3hat(i)

!Back Substitution
!Back Substitution
!Solve for NHAT-2 and NHAT-3

i=nhat-2
ai=1/ahat(i)
x1hat(i)=ai*dhat(i)
x2hat(i)=ai*d2hat(i)
x3hat(i)=ai*d3hat(i)
i=nhat-3
ai=1/ahat(i)
x1hat(i)=ai*( dhat(i) - chat(i)*x1hat(i+1) )
x2hat(i)=ai*( d2hat(i) - chat(i)*x2hat(i+1) )
x3hat(i)=ai*( d3hat(i) - chat(i)*x3hat(i+1) )
!Solve for I=NHAT-4,...,1

do i=nhat-5,1,-2
k=i+1
ai=1/ahat(k)
x1hat(k)=ai*(dhat(k)-g1hat(k)*x1hat(k+1)-g2hat(k)*x1hat(k+2))
x2hat(k)=ai*(d2hat(k)-g1hat(k)*x2hat(k+1)-g2hat(k)*x2hat(k+2))
x3hat(k)=ai*(d3hat(k)-g1hat(k)*x3hat(k+1)-g2hat(k)*x3hat(k+2))
k=i
ai=1/ahat(k)
x1hat(k)=ai*(dhat(k)-chat(k)*x1hat(k+1)-g1hat(k)*x1hat(k+2)-
$      g2hat(k)*x1hat(k+3))
x2hat(k)=ai*(d2hat(k)-chat(k)*x2hat(k+1)-g1hat(k)*x2hat(k+2)-
$      g2hat(k)*x2hat(k+3))
x3hat(k)=ai*(d3hat(k)-chat(k)*x3hat(k+1)-g1hat(k)*x3hat(k+2)-
$      g2hat(k)*x3hat(k+3))
end do

!Solve for NHAT-1 and NHAT

i=nhat-1

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aa1=v1hat*x2hat(1) + f1hat(i)*x2hat(i-2) + f2hat(i)*x2hat(i-1) +
$   ahat(i)
cc1=v1hat*x3hat(1) + f1hat(i)*x3hat(i-2) + f2hat(i)*x3hat(i-1) +
$   chat(i)
dd1=dhat(i) - v1hat*x1hat(1) - f1hat(i)*x1hat(i-2) -
$   f2hat(i)*x1hat(i-1)
i=nhat
bb2=u1hat*x2hat(1) + u2hat*x2hat(2) + f1hat(i)*x2hat(i-3) +
$   f2hat(i)*x2hat(i-2) + rhat(i)
aa2=u1hat*x3hat(1) + u2hat*x3hat(2) + f1hat(i)*x3hat(i-3) +
$   f2hat(i)*x3hat(i-2) + ahat(i)
dd2=dhat(i) - u1hat*x1hat(1) - u2hat*x1hat(2) -
$   f1hat(i)*x1hat(i-3) - f2hat(i)*x1hat(i-2)

x1=bb2/aa1
aa2=aa2 - x1*cc1
dd2=dd2 - x1*dd1
xhat(nhat)=dd2/aa2
xhat(nhat-1)=( dd1 - cc1*xhat(nhat) )/aa1

do i=1,nhat-2
    xhat(i)=x1hat(i) + xhat(nhat-1)*x2hat(i) + xhat(nhat)*x3hat(i)
end do

return
end

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